The Role of Sarcosine Metabolism in Prostate Cancer Progression


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Abstract

Metabolomic profiling of prostate cancer (PCa) progression identified markedly elevated levels of sarcosine (N-methyl glycine) in metastatic PCa and modest but significant elevation of the metabolite in PCa urine. Here, we examine the role of key enzymes associated with sarcosine metabolism in PCa progression. Consistent with our earlier report, sarcosine levels were significantly elevated in PCa urine sediments compared to controls, with a modest area under the receiver operating characteristic curve of 0.71. In addition, the expression of sarcosine biosynthetic enzyme, glycine N-methyltransferase (GNMT), was elevated in PCa tissues, while sarcosine dehydrogenase (SARDH) and pipecolic acid oxidase (PIPOX), which metabolize sarcosine, were reduced in prostate tumors. Consistent with this, GNMT promoted the oncogenic potential of prostate cells by facilitating sarcosine production, while SARDH and PIPOX reduced the oncogenic potential of prostate cells by metabolizing sarcosine. Accordingly, addition of sarcosine, but not glycine or alanine, induced invasion and intravasation in an in vivo PCa model. In contrast, GNMT knockdown or SARDH overexpression in PCa xenografts inhibited tumor growth. Taken together, these studies substantiate the role of sarcosine in PCa progression.

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Abbreviations: PCa, prostate cancer; GNMT, glycine N-methyltransferase; SARDH, sarcosine dehydrogenase; PIPOX, pipecolic acid oxidase

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Introduction
Prostate cancer (PCa) afflicts one in nine men over the age of 65 and represents the most commonly diagnosed cancer in American men [1]. Although radiation and surgery are generally effective for the majority of men, prognosis remains poor in patients with progressive disease. A distinct set of genes, proteins, and metabolites orchestrates cancer progression from precursor lesion to localized disease and finally to metastatic cancer [2]. Although gene and protein expression profiles have been extensively studied in PCa, little is known about the global metabolomic alterations. Disease-specific metabolites represent the effective end points of the molecular pathways that are initiated at genomic, transcriptomic, and proteomic levels [3] and may serve as valuable biomarkers.

In our initial study [4], we profiled approximately 1126 metabolites across 262 prostate-derived clinical samples and found that levels of sarcosine, an \( N \)-methyl derivative of the amino acid glycine, increased progressively in benign, localized PCa and metastatic disease. Subsequent studies from Jentzmik et al. confirmed elevated levels of tissue-associated sarcosine in prostate tumors compared to benign prostate [5]. In addition, our profiling study also reported elevated sarcosine levels in post-digital rectal examination (DRE) urine specimens (sediment and supernatant) from biopsy-positive PCa patients compared to biopsy-negative controls, with an area under curve (AUC) of 0.67. On the basis of this modest discriminatory power, we had nominated sarcosine as a candidate for validation and as a potential marker for inclusion into a multiplex biomarker panel for PCa detection. Since then, multiple cross-institutional validation studies to evaluate the discriminatory power of urinary sarcosine have been carried out [6–8]. Importantly, the three major studies that have used at least 50 or more samples in their validation efforts reported identical AUC for sarcosine in the range of 0.65 to 0.68 confirming our published results [6–8]. In all of these cases, sarcosine performs modestly better than serum prostate-specific antigen (PSA) in detecting biopsy-positive PCa in a clinically challenging PSA range of 4 to 10 ng/ml, while the free/total PSA ratio (%fPSA), when used as a measure, was shown to be a better predictor in the European study [9]. Furthermore, when used in a multiplex setting, sarcosine improved the overall AUC of a multiplex biomarker panel [7], consistent with our initial prediction. A recent study in individuals with serum PSA levels < 4 ng/ml demonstrated a higher predictive value for serum-associated sarcosine in detecting PCa compared to total PSA or %fPSA [10]. Within this nested PSA-controlled subset, serum sarcosine levels were significantly higher in patients with high-grade tumors. This diagnostic ability for sarcosine was also evident in yet another independent study that reported significantly elevated levels of the metabolite in presurgical urine specimens of PCa patients who developed early biochemical recurrence [11]. Together, these findings substantiate our published data describing sarcosine as an oncometabolite in PCa [4].

An overview of the sarcosine pathway is shown in Figure 1A. The major enzymes regulating sarcosine metabolism are glycine \( N \)-methyltransferase (GNMT), sarcosine dehydrogenase (SARDH), and \( L \)-pipecolic acid oxidase (PIPOX). In cells, sarcosine (\( N \)-methyl glycine) is generated by the enzymatic transfer of a methyl group from \( S \)-adenosylmethionine (SAM) to glycine with the concomitant production of \( S \)-adenosylhomocysteine (SAH). This reaction is catalyzed by the enzyme GNMT that is expressed at high levels in mammalian

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Levels of sarcosine and its pathway enzymes in PCa progression. (A) Schematic representation of the sarcosine pathway and its potential role in PCa. (B) Receiver operating characteristic curve for sarcosine in the 345 urine sediments from 211 biopsy-positive and 134 biopsy-negative individuals. Sarcosine has an AUC of 0.71. (C) Box plot showing significantly \((P < .0001)\) higher levels of sarcosine relative to alanine in urine sediments from biopsy-proven PCa and prostate biopsy-negative controls. (D) Box plot showing progressive elevation of sarcosine during progression from benign to localized (PCa) to MET samples (benign vs PCa, \( P = .0006 \) and benign vs METS, \( P < .00001 \)).
liver, exocrine pancreas, and prostate [12,13] (Figure 1A). The sarcosine-metabolizing enzymes, SARDH and PIPOX, catalyze the oxidative demethylation of sarcosine converting it back to glycine [14,15] (Figure 1A). Using both in vitro and in vivo preclinical models, this study validates sarcosine as an oncometabolite in PCa.

Materials and Methods

Clinical Specimens and Processing

Benign prostate and localized PCa tissues were obtained from a radical prostatectomy series at the University of Michigan Hospitals; metastatic PCa biospecimens (MET) were retrieved from the Rapid Autopsy Program, both of which are part of the University of Michigan Prostate Cancer Specialized Program of Research Excellence Tissue Core. Samples were collected with informed consent and prior Institutional Review Board approval at the University of Michigan (Ann Arbor, MI). Urine specimens were collected from patients screened in the urology clinic for PCa. The median age of these patients was 62.57 ± 8.14, who had clinically significant levels of PSA (8.59 ± 6.30). Urine was collected post-DRE and before biopsy. In addition, urine from patients undergoing prostatectomy, collected before surgery, was used as a positive control. In total, 652 urine specimens were examined in this study, which included 168 biopsy-negative, 261 biopsy-positive, and 210 radical retropubic prostatectomy (RRP) specimens. Six specimens from individuals with high-grade prostatic intraepithelial neoplasia (HGPIN) and seven samples from individuals with undetermined diagnosis were excluded from the study during data analysis. Further, as described later, urine samples with an alanine ratio greater than 2.5 were also excluded from the analysis. Thus, overall 507 urine specimens were used to examine the ability of sarcosine to distinguish PCa from controls at biopsy.

Frozen tissue, cell line pellets, or urine sediments were homogenized in methanol after spiking with labeled internal standards (D3-sarcosine for tissues and cell lines and D3-alanine and D3-sarcosine for urine sediments) and extracted overnight at 4°C, with constant shaking. The extraction was carried using 1:1 molar ratio of water/chloroform at room temperature. The aqueous methanolic layer was collected and dried completely. The dried methanolic extract was azoetroped twice by adding 100 μl of dimethylformamide, mixed by vortexing, and dried using a Speed Vac for 30 minutes. One hundred μl of dimethylformamide and N-methyl-N-tert-butylmethylyl trifluoroacetamide + 1% tert-butyl-dimethylchlorosilane were then added to the dried sample, capped, and incubated at 60°C for 1 hour. Following this, the sample was resuspended in ethyl acetate and injected into a gas chromatograph–mass spectrometer. Selective ion monitoring was used for quantification. Importantly, both these metabolites were baseline separated before their quantification. The amount of sarcosine and alanine in the sample was calculated by measuring the peak area of the native sarcosine and alanine (both having m/z = 232) to that corresponding to spiked isotope-labeled sarcosine and alanine (both m/z = 235; Figure W1, A and B), as described in our previous study [4]. Following this, for tissues, the level of sarcosine was normalized to the tissue weight. In case of urine sediments, a ratio of normalized sarcosine to normalized alanine was derived. Urine specimens having normalized alanine levels (m/z = 232/235) > 2.5-fold were removed and not used for further analysis. This resulted in a total of 134 biopsy-negative, 211 biopsy-positive, and 162 RRP samples that were used to evaluate the biomarker potential of sarcosine. To evaluate the accuracy and limit/linearity of detection of sarcosine, we performed replicate experiments (technical and biological) as well as developed a calibration curve for the metabolite in urine sediments (Figure W2, A and B, calibration curve for alanine and sarcosine and Figure W2, C and D, technical and biologic replicate measurements).

Plasmids, Cell Lines, and Cell Culture

Mammalian expression construct of SARDH and PIPOX were generated by subcloning the polymerase chain reaction (PCR) product into the pEF vector using NotI and XbaI sites. Full-length GNMT was cloned in lentiviral vector backbone FUGW (kindly gifted by David Baltimore Laboratory, Pasadena, CA) and virus generated for further transduction. The primers used are listed in Table W1. The benign immortalized prostate cell line RWPE and PCa cell lines DU145, PC3, and LNCaP were obtained from the American Type Culture Collection (ATCC, Manassas, VA). DU145, LNCaP, and PC3 PCa cells were cultured in RPMI supplemented with 10% FBS (Invitrogen, Carlsbad, CA). RWPE cells were grown in keratinocyte serum-free medium (Invitrogen) supplemented with 5 ng/ml human recombinant epidermal growth factor and 0.05 mg/ml bovine pituitary extract. All the cells used in this study were tested for mycoplasma using a PCR-based method (Lonza, Norwalk, CT). For stable knockdown of GNMT, SARDH, or PIPOX, human lentiviral shRNA clones targeting the respective gene or nonsilencing (control) lentiviral shRNA in pLKO puromycin vectors were purchased from Sigma (St Louis, MO). Viral particle infections were carried out in the presence of polybrene (8 μg/ml) in 50% to 60% confluent DU145 and RWPE cells. After 48 hours, transduced cells were grown in DU145 and RWPE culture media containing puromycin (2 μg/ml). Three weeks later, stable cells were plated into 96-well plate for clonal selection. GNMT, SARDH, and PIPOX knockdown were confirmed in pooled and single clones by quantitative PCR (qPCR) and individual clones showing highest levels of knockdown were further expanded. For overexpression studies, stable cell lines were created by transfecting the FLAG-SARDH and FLAG-PIPOX construct and vector control into DU145 cells using FuGENE 6 (Roche, Indianapolis, IN) and resulting clones were selected using 300 μg/ml G418 (Invitrogen).

Immunoblot Analysis and Antibodies

Protein extracts were prepared from benign prostate, localized PCa, and metastatic tumor tissues, and standard immunoblot analysis was performed. Briefly, prostate tissues were homogenized in NP-40 lysis buffer containing 50 mM Tris-HCl (pH 7.4), 1% NP-40 (Sigma), and complete protease inhibitor cocktail (Roche). Fifteen micrograms of protein extract was mixed with sodium dodecyl sulfate sample buffer and run on a 4% to 12% Bis-Tris gel under reducing conditions. The separated proteins were transferred onto polyvinylidene fluoride membranes (Amersham Pharmacia Biotech, Piscataway, NJ) and incubated for 1 hour in blocking buffer (Tris-buffered saline with 0.1% Tween and 5% nonfat dry milk), followed by incubation with the appropriate primary antibodies overnight at 4°C. After washing with Tris-buffered saline with 0.1% Tween buffer, the membrane was incubated with HRP-linked donkey, anti-rabbit IgG antibody (GE Healthcare Life Sciences, Piscataway, NJ) at 1:5000 for 1 hour at room temperature. Specific signals were visualized by using ECL Plus Western Detection System (GE Healthcare Life Sciences). The anti-FLAG (M2) antibody was obtained from Sigma-Aldrich (St Louis, MO). Antibody against GNMT (H-225) was acquired from Santa Cruz Biotechnology (Santa Cruz, CA). SARDH and PIPOX antibodies were obtained from Abnova (Taipei, Taiwan), respectively. Antibody for poly(ADP-ribose) polymerase (PARP) was purchased...
from Cell Signaling Technology (Danvers, MA). The anti-β-actin antibody was purchased from Abcam (Cambridge, MA). Protein levels for actin were used as loading control.

**Quantitative PCR**

Total RNA was isolated from DU145-shNS−, DU145-shGNMT−, RWPE-shNS−, RWPE-shSARDH−, or RWPE-shPIPOX−-transduced cells (Qiagen, Gaithersburg, MD). qPCR was performed using SYBR Green dye on the StepOne Real-Time PCR System (Applied Biosystems, Foster City, CA) [16]. Briefly, 1 μg of total RNA was reverse transcribed into cDNA using SuperScript III (Invitrogen) in the presence of random primers (Invitrogen). All reactions were performed in triplicates with SYBR Green Master Mix (Applied Biosystems) and 25 ng of both the forward and reverse primers according to the manufacturer’s recommended thermocycling conditions and then subjected to melt curve analysis. Threshold levels for each experiment were set during the exponential phase of the qPCR reaction using Sequence Detection Software version 1.2.2 (Applied Biosystems). The relative quantity of the target gene was computed for each sample using the ΔΔCt method by comparing mean Ct of the gene to the mean Ct of the housekeeping gene, glyceraldehyde 3-phosphate dehydrogenase (GAPDH) [17]. All oligonucleotide primers were synthesized by Integrated DNA Technologies (Coralville, IA). Sequences of all primers used in qPCR analyses are listed in Table W1.

**Flow Cytometry Analysis**

shGNMT-DU145 cells were collected and fixed with 5 ml of ice-cold methanol for 1 hour at 4°C. Fixed cells were resuspended in phosphate-buffered saline solution containing RNase and propidium iodide (200 μg/ml; Sigma), and flow cytometry analysis was performed. Cell cycle profiles were obtained using a BD Biosciences FACSCalibur flow cytometer and CellQuest software. Apoptosis was assessed using sub-G0/G1 DNA content.

**Immunohistochemistry**

A study pathologist (R.M.) reviewed the slides of all tissue samples from the radical prostatectomy series and designated areas of interest. These slides were used as a template for tissue microarray (TMA) construction. All TMAs were assembled using the manual tissue arrayer (Beecher Instruments, Silver Spring, MD) with at least three tissue cores sampled from each donor block. Histologic diagnosis of the tissue cores was verified by standard hematoxylin and eosin staining of the initial paraffin block. These slides were used as a template for tissue microarray (TMA) construction. All TMAs were assembled using the manual tissue arrayer (Beecher Instruments, Silver Spring, MD) with at least three tissue cores sampled from each donor block. Histologic diagnosis of the tissue cores was verified by standard hematoxylin and eosin staining of the initial paraffin block. These slides were used as a template for tissue microarray (TMA) construction. All TMAs were assembled using the manual tissue arrayer (Beecher Instruments, Silver Spring, MD) with at least three tissue cores sampled from each donor block.
and implanted subcutaneously into the left flank regions of the mice. Growth in tumor volume was recorded weekly using digital calipers, and tumor volumes were calculated using the formula \( \frac{\pi}{6}(L \times W^2) \), where \( L \) is the length of tumor and \( W \) is the width. Tumor growth inhibition was defined as percent decrease in the mean tumor volume for shGNMT relative to shNS vector and SARDH over-expressing cells relative to the vector control. All experimental procedures involving mice were approved by the University Committee on Use and Care of Animals at the University of Michigan and conform to their relevant regulatory standards.

### Statistical Analysis

To examine the biomarker potential of sarcosine for PCa, a classification model based on logistic regression was developed. Here, the value of normalized sarcosine was used as the predictor variable, and biopsy status, i.e., positive or negative, was used as the binary outcome variable. The performance of the model was examined by using the area under the receiver operating characteristic curve (AUROC). AUROC was determined using a cross-validation approach, by partitioning the specimens \( n = 345 \), both biopsy positive and negative equally into training and testing groups. The cross-validation was carried out by repeating this partitioning process 5000 times. The receiver operating characteristic reflecting the mean AUROC across all the iterations along with the associated 95% confidence band is shown in Figure 1B.

For the functional studies, all values are presented as means ± SEM. The significant differences between the groups were analyzed by a Student’s \( t \) test and a \( P \) value < .05 was considered significant.

### Results

#### Sarcosine Levels Are Significantly Elevated in PCa

We reconfirmed our earlier results on the modest ability of sarcosine to detect PCa at biopsy, using a training/testing approach, on independent sets of urine specimens. To build the training model, we used 211 biopsy-positive and 134 biopsy-negative urine sediments, and implanted subcutaneously into the left flank regions of the mice. Growth in tumor volume was recorded weekly using digital calipers, and tumor volumes were calculated using the formula \( \frac{\pi}{6}(L \times W^2) \), where \( L \) is the length of tumor and \( W \) is the width. Tumor growth inhibition was defined as percent decrease in the mean tumor volume for shGNMT relative to shNS vector and SARDH over-expressing cells relative to the vector control. All experimental procedures involving mice were approved by the University Committee on Use and Care of Animals at the University of Michigan and conform to their relevant regulatory standards.

Figure 2. GNMT, SARDH, and PIPOX mRNA and protein expression are dysregulated in PCa. (A) Quantitative SYBR Green reverse transcription–PCR (qRT-PCR) assessment of GNMT, SARDH, and PIPOX expression in prostate tumor specimens. Expression was determined in a cohort of 12 benign adjacent prostate (benign), 14 localized PCa, and 14 metastatic PCa (MET) tissues. For qRT-PCR, expression of target genes was normalized to the expression of the housekeeping gene, GAPDH. (B) Immunoblot analysis showing GNMT, SARDH, and PIPOX expression in benign, PCA, and MET. The protein levels of actin were used as loading control. (C) Representative prostate tissue sections stained with an antibody to GNMT. (Left) GNMT expression in benign tissue was observed primarily in cytoplasm (green arrow). (Right) In clinically localized PCa, the expression of GNMT was increased in the cytoplasm (red arrow) compared to weak expression in benign gland from the same sample (green arrow). (D) Box plot of GNMT product score across PCa progression.
Figure 3. The role of the sarcosine-generating enzyme, GNMT, in prostate cell lines. (A) Benign immortalized prostate RWPE cells were transduced with GFP-GNMT lentivirus or mock or GFP control. Increased levels of sarcosine were found in GFP-GNMT lentivirus–transduced RWPE cells compared to mock or GFP control. GNMT overexpression also increased invasion in RWPE cells as measured by Boyden chamber Matrigel invasion assay. Representative photomicrographs showing cell invasion assay (top inset) are shown. (B) Cell proliferation assay using pooled shGNMT, shGNMT clone 7, shGNMT clone 9, or shNS cells at the indicated time points showing decreased cell proliferation in shGNMT knockdown cells. (C) shGNMT clone 7 and shGNMT clone 9 or shNS DU14 cells were plated for 48 hours. The amount of sub-G0/G1 cells was calculated using the CellQuest program for fluorescence-activated cell sorting (FACS). Cleavage of PARP by immunoblot assessment is shown in the top inset. (D) Assessment of sarcosine levels by GC-MS showing decreased levels of sarcosine in shGNMT knockdown cells compared to shNS vector. The invasion was quantitated by absorbance and cells were photographed after invasion through Matrigel and stained with crystal violet (top inset). (E) Pooled shGNMT, shGNMT clone 7, or shGNMT clone 9 blocked anchorage-independent growth in soft agar assay compared to shNS vector control. Representative photomicrographs showing soft agar (top inset) are shown. All experiments were independently performed in triplicates. (F) Exogenous sarcosine or alanine was spiked to shGNMT clone 7 or shGNMT clone 9 cells and invasion assay was performed. The addition of sarcosine resulted in the rescue of invasion phenotype in these GNMT knockdown cells. Data shown in the figure represent means ± SEM. Asterisks indicate significant comparisons (\( P < .05 \), two-sided Student’s t test).
collected post-DRE and pre-biopsy. Using a logistic regression model, sarcosine was found to have a cross-validated AUROC of 0.71, for distinguishing biopsy-positive patients from biopsy-negative controls (Figure 1B), verifying our earlier findings [4]. Sarcosine levels were found to be significantly higher in urine sediments ($P < .0001$; Figure 1C) derived from biopsy-positive PCa patients compared to biopsy-negative controls. In addition, using a similar approach, the AUROC for sarcosine to distinguish biopsy-negative controls from the combined group consisting of biopsy-proven PCa as well as patients who underwent RRP was 0.695 (Figure W3). Following this, we next verified elevated sarcosine levels in localized PCa and metastatic tissues. Sarcosine levels were significantly elevated in localized PCa ($n = 11$) compared to benign adjacent prostate samples ($n = 11$, $P = .0006$; Figure 1D). The levels of the metabolite were even greater in metastatic samples ($n = 10$) compared to localized PCa ($P < .00001$; Figure 1D). Interestingly, during the course of these validations, we observed that the metabolite levels in noncancer-containing tissues were sensitive to the “field effect” generated by the adjacent tumor. Thus, as shown in Figure W4, the levels of sarcosine were elevated and comparable in both the site of metastatic tumor and the adjacent pathologically normal tissue. The levels of the metabolite, however, were undetectable in the benign tissues that were distant from the metastatic site in the same patient (Figure W4).

**Transcript and Protein Levels of GNMT, SARDH, and PIPOX Are Dysregulated during PCa Progression**

Having confirmed elevated levels of sarcosine in localized/metastatic tissues as well as biopsy-positive urine sediments, we examined the role of sarcosine in PCa development and progression. To this end, we first measured the mRNA levels of the enzymes in the sarcosine pathway that included GNMT, SARDH, and PIPOX in prostate tissues. We carried out SYBR Green real-time qPCR on 12 benign prostates, 14 localized PCa, and 14 metastatic samples. Our results confirmed
earlier results allude to a stepwise increase in levels of sarcosine, from benign prostate tissue to localized PCa to metastatic disease, as reported previously. To confirm this, we transduced these benign cells with a lentiviral construct expressing GNMT [or green fluorescent protein (GFP) as control] and measured their sarcosine levels as well as examined its effect on proliferation and invasive potential. Overexpression of GNMT in RWPE cells elevated sarcosine levels (Figure 3A) with concomitant increase in invasion (Figure 3A; P = .0009) but had no effect on cell proliferation (Figure W5).

Subsequently, we carried out a stable knockdown of the enzyme in DU145 PCa cells that are known to be invasive. Here, we used two independent GNMT-specific shRNA (shGNMT), individually as well as in a pooled format, along with nontargeted shRNA (shNS) as a control. In all of these instances, we observed >70% reduction in GNMT expression both at the mRNA (Figure W6A) and protein levels (Figure W6B). This was concomitantly associated with a reduction in cell proliferation compared to nontarget-transduced controls (54% and 60% reduction, respectively; Figure 3B). This was reflected in flow cytometry data, which showed an increase in the sub-G1 fraction, in the knockdown cells (Figure 3C). Furthermore, these cells also exhibited a greater percentage of cell death, possibly due to apoptosis. To confirm this, we evaluated PARP family of proteins that are known to be cleaved during programmed cell death. As expected, cells in which GNMT was knocked down showed higher levels of cleaved PARP compared to controls (Figure 3C, top inset), suggestive of apoptosis, as an additional mechanism leading to reduced cell numbers.

In addition to changes in cell numbers, GNMT knockdown cells also showed >65% reduction in both invasion and anchorage-dependent growth compared to nontarget-transduced controls, as assessed by Boyden chamber Matrigel assay and soft agar colony-forming assay (Figure 3, D, gray bars, and E; P < .005 for both invasion and soft agar assay). Notably, all of the above phenotypes in cells with reduced GNMT expression were also associated with markedly lower levels of sarcosine (Figure 3D, black bars). Consistent with this, addition of sarcosine to these cells partially rescued their invasive properties, while addition of a control metabolite alanine failed to rescue the invasive phenotype (Figure 3F; P < .05). Addition of exogenous sarcosine did not completely rescue invasive phenotype of DU145 cells expressing low levels of GNMT possibly because endogenous SARDH and PIPOX activity converted sarcosine to glycine. This needs to be verified by flux analysis using isotopically labeled sarcosine and the study of the kinetics of 13C incorporation into the breakdown products. Furthermore, we believe that wild-type DU145 cells are highly invasive, and hence, addition of sarcosine to these cells may not show a significant increase in invasion. Similar results were also obtained with LNCaP and PC3 (both PCa) cell lines harboring GNMT knockdown but not RWPE (benign) cells (Figures W7–W9), reinforcing the importance of the metabolite in PCa.

**Figure 5.** The role of sarcosine in PCa growth in vivo. (A) CAM invasion assay performed using RWPE cells transduced with red fluorescent protein virus for visualization. After puromycin selection, cells that were treated with vehicle, sarcosine, glycine, or alanine as indicated. Seventy-two hours after implantation, the upper CAM was harvested. Frozen sections were created and stained for hematoxylin and eosin (left column), human-specific cytokeratin (immunohistochemistry, middle column), or chicken-specific type IV collagen (green immunofluorescence, right column). Arrowheads indicate cells that invaded through the upper CAM. Representative images are shown. Scale bars, 200 μM. (B) CAM intravasation assay performed using RWPE cells treated with vehicle, sarcosine, glycine, or alanine as indicated. Seventy-two hours after implantation, the lower CAM was harvested. Total DNA was isolated from the lower CAM, and qPCR was performed using human-specific Alu PCR primers. Total cell number was determined by comparing to a standard curve derived by using known quantities of input RWPE cells.

**Perturbation of Sarcosine-Generating Enzyme, GNMT, Modulates Neoplastic Progression of Prostate Cell Lines**

Our data thus far suggest an elevation of sarcosine metabolism in PCa progression. Furthermore, our earlier results that were based on transient knockdown of enzymes in the sarcosine pathway indicated a tumor-promoting role for this metabolite in PCa. To explore this further, we examined the effect of ectopic overexpression of GNMT in benign prostate epithelial cells, RWPE. To achieve this, we transduced these benign cells with a lentiviral construct expressing GNMT [or green fluorescent protein (GFP) as control] and measured their sarcosine levels as well as examined its effect on proliferation and invasive potential. Overexpression of GNMT in RWPE cells elevated sarcosine levels (Figure 3A) with concomitant increase in invasion (Figure 3A; P = .0009) but had no effect on cell proliferation (Figure W5).

A Functional Role for SARDH and PIPOX in PCa Progression In Vitro

Next, we examined the effect of perturbing the enzyme machinery that regulates the metabolism of sarcosine to glycine on the phenotype of the prostate cell lines. As described earlier in Figure 1A, SARDH...
and PIPOX convert sarcosine back to glycine. First, we examined the effect of SARDH and PIPOX overexpression on growth and invasive potential of DU145 PCa cells (Figure W10). Notably, in case of DU145 cells, overexpression of these enzymes markedly decreased sarcosine levels and attenuated invasion (Figure 4A), while only SARDH overexpression reduced proliferation (Figure 4B). Consistent with this, only overexpression of SARDH but not PIPOX reduced colony formation in soft agar relative vector control (Figure 4C). In contrast, knockdown of SARDH in RWPE cells increased sarcosine levels and resulted in increased proliferation (Figure 4D), invasion (Figure 4E),

Figure 6. The role of sarcosine pathway enzymes, GNMT and SARDH, in PCa growth in vivo. (A) CAM intravasation assay was performed using shGNMT clone 7, SARDH clone 5, and vector control in DU145 cells. Number of intravasated DU145 cells and tumor weight was decreased in both shGNMT and SARDH overexpressing xenografted cell lines compared to vector control. (B) Liver metastasis in chicken embryos was assessed 8 days following implantation of either shGNMT clone 7 or SARDH clone 5 or vector control cells onto the upper CAM. Total number of metastasized cells were quantified and found to be significantly decreased in both shGNMT and SARDH overexpressing xenografted cell lines compared to vector control. (C) shGNMT (clones 7 and 9) decreased DU145 tumor growth in mice. Means ± SEM are shown, *P < .05. (D) Stably overexpressed SARDH (pool and clone 5) decreased DU145 tumor growth in mice. Means ± SEM are shown, *P < .05. (E) Box plots showing decreased sarcosine levels in shGNMT (clones 7 and 9) mouse xenograft compared to shNS vector control. (F) Box plot showing decreased sarcosine levels in SARDH overexpressed (pool and clone 5) mouse xenograft tumors compared to vector control.
and anchorage-independent growth (Figure 4F). However, knockdown of PIPOX in RWPE increased sarcosine levels and made them invasive, while having no effect on their proliferation and anchorage-independent growth.

**Sarcosine-mediated Invasion and Intravasation In Vivo**

Further developing our *in vitro* findings, we examined the ability of sarcosine, glycine, and alanine to trigger local invasion and intravasation, in nontumorigenic RWPE cells, using a CAM assay [23]. Consistent with our *in vitro* data, addition of sarcosine but not glycine or alanine induced invasion and intravasation in RWPE cells that migrated across the upper CAM junction (Figure 5, A and B, arrows; *P* < .01). GNMT knockdown or SARDH overexpression in DU145 PCa cells, however, resulted in a significant reduction in tumor weight (*P* = .02 and *P* = .03, respectively) as well as decreased intravasation (*P* = .01 for GNMT knockdown and *P* = .05 for SARDH overexpression, respectively; Figure 6A), and ability of the cells to metastasize to the liver (*P* < .05), compared to controls (Figure 6B). Similar results were obtained using PC3 PCa cells as shown in Figure W11.

To confirm these *in vivo* results further, we examined the effect of GNMT knockdown and SARDH overexpression on *in vivo* tumor growth using a mouse xenograft model. We implanted DU145 cells containing either stable GNMT knockdown or ectopic overexpression of SARDH and their respective controls in nude male mice. At 5 weeks post-implantation, DU145 cells containing GNMT knockdown formed significantly smaller tumors (~75% reduction, *P* = .0001) compared to control cells (Figure 6C). Similarly, SARDH overexpressing tumors were also significantly reduced in size (~70% reduction, *P* = .0001) compared to their corresponding controls (Figure 6D). As expected, in both GNMT knockdown and SARDH overexpressing tumors, sarcosine levels were significantly reduced compared to controls (Figure 6, E and F). Furthermore, the tumors containing GNMT knockdown also had significantly elevated levels of SAM and lower levels of SAH, indicative of reduced activity through the GNMT-catalyzed reactome (Figure W12, A and B; *P* = .05).

**Discussion**

Our prior unbiased survey of the PCa metabolome identified sarcosine as a potential marker for aggressive disease that could be detected noninvasively in urine [4]. Further, *in vitro* studies showed that exogenous addition of sarcosine induced an invasive phenotype in benign prostate epithelial cells [4]. In the present study, we confirmed elevated levels of sarcosine in PCa tissues and urine and establish a role for sarcosine in promoting PCa growth and progression, using both *in vitro* and *in vivo* models of the disease.

Our studies on tissues revealed that the levels of sarcosine in pathologically benign tissues could potentially be influenced by the presence of an adjacent tumor. PCa like many tumors is known to exhibit a “field effect” that influences the metabolism of adjacent benign tissues [4,25]. Although the sensitivity of sarcosine levels to the underlying “field effect” needs to be more extensively characterized, it could partially explain the modest change in the metabolite levels seen in organ-confined PCa compared to its matched benign adjacent tissues [4]. Irrespective of this, our data confirm SARDH as a potential tumor suppressor in PCa, consistent with a recent report by Chen et al. [29]. In summary, this study reiterates an important role for sarcosine in PCa progression and highlights the potential of using the metabolite to develop a multiplex urine-based marker for PCa detection.

**Acknowledgments**

We thank V. Vitvitsky from R. Banerjee Laboratory for SAM and SAHA estimations and X. Cao, Robert J. Lonigro, and N. Palanisamy for helpful discussions. We also thank Khalid Suleman and Rong Zhao for technical support.

**References**


Supplemental Reference

**Figure W1.** (A) Extracted ion chromatogram of sarcosine and alanine in urine. (B) Selected ion monitoring of alanine and sarcosine.
Table W1. Sequence of Gene-Specific PCR Primers Used for Cloning and qPCR.

<table>
<thead>
<tr>
<th>Gene-Specific Primer</th>
<th>Primer Sequence</th>
</tr>
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<tbody>
<tr>
<td>XhoI-GNMT F</td>
<td>5′-TTTTCCCTTTTTGCCGCCGCTgggacagggacggacc-3′</td>
</tr>
<tr>
<td>EcoR1-GNMT R</td>
<td>5′-CCGGAAATTCCctttccctttctttgcacc-3′</td>
</tr>
<tr>
<td>Nor1-SARDH F</td>
<td>5′-TTTTTCTTTTTGCCGCCGCTcacaagggccgacc-3′</td>
</tr>
<tr>
<td>XhoI-SARDH R</td>
<td>5′-CTAGTCTAGAtcagtagattcccttcaccct-3′</td>
</tr>
<tr>
<td>Nor1-PIPOX F</td>
<td>5′-TTTTTCTTTTTGCCGCCGCTgacgcaagaccct-3′</td>
</tr>
<tr>
<td>XhoI-PIPOX R</td>
<td>5′-CTAGTCTAGAcgaaggggtggtggtggtggtg-3′</td>
</tr>
<tr>
<td>GNMT F*</td>
<td>5′-CTTCATCCACGGCTGTCCTAGA-3′</td>
</tr>
<tr>
<td>GNMT R*</td>
<td>5′-TCCCCCATCTTCCAGACAGACG-3′</td>
</tr>
<tr>
<td>SARDH F*</td>
<td>5′-CTAGTAGATGTCAGACCCT-3′</td>
</tr>
<tr>
<td>SARDH R*</td>
<td>5′-GTTCATCTATGACCTGTGCTG-3′</td>
</tr>
<tr>
<td>PIPOX F*</td>
<td>5′-CTTCATCTTTCCTTGACCTT-3′</td>
</tr>
<tr>
<td>PIPOX R*</td>
<td>5′-GAAGGGACACAGATACCCTGTC-3′</td>
</tr>
</tbody>
</table>

F and R stand for forward and reverse primers respectively.

*Denotes set of primers used for qPCR analysis.
Figure W4. Sarcosine levels in metastatic and benign tissues of five different cancer patients. Sarcosine levels were found to be comparably high in both metastatic tumor (black) and its matched adjacent pathologically benign tissues (gray), while still being undetectable in distal nontumor tissues resected from the same patient [1].

Figure W3. AUROC for sarcosine in the 652 urine sediments from 168 biopsy-negative, 261 biopsy-positive, and 210 RRP individuals was 0.695.
Figure W5. GNMT overexpression does not affect proliferation of benign RWPE prostatic epithelial cells. The benign immortalized prostate cell line RWPE was transduced for overexpression of GNMT or GFP vector control. We observed no significant difference in cell proliferation in RWPE-GNMT overexpressing cells relative to either GFP transduced or mock.

Figure W6. qRT-PCR and immunoblot analysis of GNMT knockdown in DU145 cells. (A) qPCR analysis of GNMT transcript in stable shGNMT knockdown cells (stably selected pool and clones). (B) Western blot analysis showing stably puromycin-selected clones of GNMT knockdown in DU145 cell model.
Figure W7. Validation of GNMT knockdown followed by assessment of sarcosine levels in RWPE, PC3, and LNCaP cells. qPCR analysis of GNMT transcript in stable shGNMT knockdown (A) RWPE, (B) PC3, and (C) LNCaP cells. Sarcosine levels were assessed by GC-MS in stable shGNMT knockdown (D) RWPE, (E) PC3, and (F) LNCaP cells. We observed significant decrease in sarcosine levels after stable GNMT knockdown in PC3 and LNCaP cells.
Figure W8. GNMT knockdown decreases cell proliferation in (A) PC3 and (B) LNCaP cells. GNMT knockdown in metastatic prostate carcinoma (PC3) and androgen-responsive (LNCaP) cells resulted in the significant decrease in cell proliferation.

Figure W9. GNMT knockdown attenuates invasion in PC3 and LNCaP cells. (A) PC3 and (B) LNCaP cells were transduced with shGNMT, or shNS vector control, stable cells were generated and assayed for cell invasion.
Figure W10. Validation of overexpression and knockdown of SARDH and PIPOX in DU145 and RWPE cells, respectively. (A) Western blot showing stably transfected FLAG-tagged PIPOX and FLAG-tagged SARDH expression (pool and clones; DU145 cells). Actin was used as loading control. qPCR analysis of RNA isolated in parallel to the invasion assay of (B) SARDH and (C) PIPOX.
Figure W11. Tumor weight decreases in GNMT knockdown PC3 xenografts in CAM assay compared to vector control.

Figure W12. Assessment of SAM and SAH in GNMT knockdown mouse xenograft. (A) Box plot of SAM. (B) Box plot of SAH.